| | (FILE ROME ENTERED AT 14:09:36 ON 02 NOV 2005) |
|-----|--|
| | FILE 'REGISTRY' ENTERED AT 14:09:50 ON 02 NOV 2005 |
| L1 | FILE 'CAPLUS' ENTERED AT 14:10:09 ON 02 NOV 2005 STRUCTURE UPLOADED S L1 |
| L2 | FILE 'REGISTRY' ENTERED AT 14:10:42 ON 02 NOV 2005 10 S L1 FULL |
| | FILE 'CAPLUS' ENTERED AT 14:10:45 ON 02 NOV 2005 |
| L3 | 11 S L2 FULL |
| L4 | 7 S L3 AND PY<2003 |
| L5 | 0 S L4 AND ORTHORHOMBIC |
| L6 | 0 S L4 AND CRYSTAL? |
| T.7 | O S IA AND DOLYMODDY |

•• •

.

.

·

4-[6-ACETYL-3-[3-(4-ACETYL-3-HYDROXY-2-PROPYLPHENYLTHIO)PROPOXY]-2-PROPYLPHENOXY]BUTYRIC ACID

=> s l1 full

REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

FULL SEARCH INITIATED 14:10:42 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 422943 TO ITERATE

100.0% PROCESSED 422943 ITERATIONS

00 00 04

SEARCH TIME: 00.00.04

L2 10 SEA SSS FUL L1

L3 11 L2

=> s 13 and py<2003 22790221 PY<2003

7 L3 AND PY<2003

=> s 14 and orthorhombic

59874 ORTHORHOMBIC

L5 0 L4 AND ORTHORHOMBIC

=> s l4 and crystal?

1695429 CRYSTAL?

L6 0 L4 AND CRYSTAL?

=> d l4 1-7 ibib abs hitstr

L4 ANSWER 1 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2001:564826 CAPLUS

DOCUMENT NUMBER:

135:142249

TITLE:

Eye drop compositions containing leukotriene

10 ANSWERS

antagonist KCA-757

INVENTOR(S):

Kodaira, Hiromichi; Kozuka, Hitoshi

PATENT ASSIGNEE(S): SOURCE:

Kyorin Pharmaceutical Co., Ltd., Japan

PCT Int. Appl., 17 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent Japanese

LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | | | KIND DATE | | APPLICATION NO. | | | | | DATE | | | | | | | |
|---------------|------|-----|-----------|-----|-----------------|-----|------|------|-----|-------|-------|-------|------|-----|-----|-------|-------|
| | | | | | | | | | | | | | | | | | |
| WO 2001054684 | | | | | WO 2001-JP430 | | | | | | | | | | | | |
| | W: | | | | | | ΑZ, | | | | | | | | | | |
| | | | | | | | | | | | | | | | | | IL, |
| | | | | | | | KΡ, | | | | | | | | | | |
| | | MG, | MK, | MN, | MW, | MX, | NO, | NZ, | PL, | PT, | RO, | RU, | SD, | SE, | SG, | SI, | SK, |
| | | | | | | | UÀ, | UG, | US, | UZ, | VN, | YU, | ZA, | ZW, | AM, | ΑZ, | BY, |
| | | - | - | - | RU, | • | | | | | | | | | | | |
| | RW: | | | | | | MZ, | | | | | | | | | | |
| | | | | | | | GB, | | | | | | | | | TR, | BF, |
| | | | | | | | GΑ, | | | | | | | | | | |
| | | | | | | | | | | | | | | | | | L24 < |
| | | | | | | | | | | | | | | | | | 124 < |
| EΡ | 1250 | 924 | | | A1 | | 2002 | 1023 |] | EP 20 | 001- | 94678 | 38. | | 20 | 00101 | 124 < |
| | R: | AT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, | GR, | IT, | LI, | LU, | NL, | SE, | MC, | PT, |
| | | ΙE, | SI, | LT, | LV, | FI, | RO, | MK, | CY, | AL, | TR | | | | | • | |
| TW | 5260 | 51 | | | В | | 2003 | 0401 | 7 | FW 20 | 001-9 | 9010 | 1616 | | 20 | 0101 | L29 |

US 2003083378 **A1** 20030501 US 2002-181436 20020725 PRIORITY APPLN. INFO.: JP 2000-17403 A 20000126

WO 2001-JP430 W 20010124 Disclosed are eye drops containing a potent and selective leukotriene

AB antagonist. Specifically, stable eye drops of an aqueous solution or suspension type, containing as the active ingredient 4-[6-acetyl-3-[3-[(4-acetyl-3hydroxy-2-propylphenyl)thio]-propoxy]-2-propylphenoxy]butyric acid (KCA-757). An eye drop composition containing KCA-757 0.5 g, 0.1 M NaOH 20 mL, potassium dihydrogenphosphate 0.004, sodium hydrogenphosphate 0.089, NaCl 0.8 g, and 0.1 M HCl q.s. to pH 8.5, and water q.s. to 100 mL was formulated.

IT 125961-82-2

CN

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(eye drop compns. containing leukotriene antagonist KCA-757)

RN 125961-82-2 CAPLUS

> Butanoic acid, 4-[6-acetyl-3-[3-[(4-acetyl-3-hydroxy-2propylphenyl)thio]propoxy]-2-propylphenoxy]- (9CI) (CA INDEX NAME)

$$HO_2C-(CH_2)_3-O$$
 AC
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$

REFERENCE COUNT: 25 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 2 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1999:205557 CAPLUS

DOCUMENT NUMBER: 130:287054

TITLE: Powder inhalants containing

[(propylphenyl)thio]propoxy]propylphenoxybutyrate for

the treatment of asthma

INVENTOR (S): Hoshino, Ryoichi

PATENT ASSIGNEE(S): Kyorin Pharmaceutical Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE | | |
|-------------------------|-------|--------------|-----------------|------------|--|--|
| | | | | · · | | |
| JP 11079985 | A2 | 19990323 | JP 1997-251280 | 19970901 < | | |
| PRIORITY APPLN. INFO.: | | | JP 1997-251280 | 19970901 | | |
| AB Powder inhalants for | r tho | treatment of | | | | |

Powder inhalants for the treatment of asthma comprise powdery 4-[6-acetyl-3-[3-(4-acetyl-3-hydroxy-2-propylphenylthio)propoxy]-2propylphenoxy]butyric acid (I) as an active ingredient. I in combination with lubricants is suspended in an aqueous solution of polymers and spray dried to give a fine powder having an average particle diam $\leq 6~\mu m$. The powders show little self-cohesive properties and little adhesion to a dispersing device. Hydroxypropyl Me cellulose 1.5 g was dissolved in 380 g distilled water and to the solution 0.5 g sucrose fatty acid ester was added, followed by 18 g I. The dispersion was subjected to a high-pressure homogenization and spray-drying to give a dry powder inhalant. IT125961-82-2

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (manufacture of antiasthmatic powder inhalants containing

[(propylphenyl)thiopropoxy]propylphenoxybutyrate and polymers and lubricants)

RN 125961-82-2 CAPLUS

$$HO_2C-(CH_2)_3-O$$
 AC
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$

ANSWER 3 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1996:379374 CAPLUS

DOCUMENT NUMBER: 125:58104

TITLE: Preparation of phenoxycarboxylic acid derivatives as

antiallergy agents

INVENTOR(S): Matsumoto, Toyomi; Ishiguro, Juji; Myashita, Kunio;

Kitamura, Genichi

PATENT ASSIGNEE(S): Kyorin Seiyaku Kk, Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 4 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1.

PATENT INFORMATION:

CN

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|------|----------|-----------------|------------|
| | | | | |
| JP 08081412 | A2 | 19960326 | JP 1994-244636 | 19940913 < |
| PRIORITY APPLN. INFO.: | | | JP 1994-244636 | 19940913 |
| A | | | | |

OTHER SOURCE(S): CASREACT 125:58104; MARPAT 125:58104

Ι

Y (CH₂)
$$_{m}$$
X — COMe

Pr O (CH₂) $_{n}$ CO₂H

$$Ac$$
 X^1CONMe_2 HO Pr

ΙI

Ac
$$\longrightarrow$$
 X1 (CH₂) mX \longrightarrow COMe
HO Pr Pr O (CH₂) nCO₂H IV

AB The title derivs. IV (m = 2-5; n = 3-8; X1 = S, 0; X = O, S, SO, SO2; X1 = X ≠ O), useful as antiallergy agents (no data), are prepared by treating phenoxycarboxylic acids I (Y = halo) with hydroxybenzenes III, which is formed by hydrolysis of hydroxyphenyl carbamates II, in one pot. A mixture of 10 g S-(4-acetyl-3-hydroxy-2-propylphenyl) N,N-dimethylthiocarbamate and KOH in H2O was treated at 95° for 1.5 h, then treated with 12.7 g 4-[6-acetyl-3-hydroxy-3-(3-chloropropoxy)-2-propylphenoxy]butyric acid at 35-40° for 21 h to give 15.2g 4-[6-acetyl-3-(3-(4-acetyl-3-hydroxy-2-propylphentylthio)propoxy)-2-

propylphenoxy]butyric acid.

IT 125961-82-2P

RN

CN

RL: IMF (Industrial manufacture); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of phenoxycarboxylic acid as antiallergy agent from phenoxycarboxylate and hydroxyphenyl carbamate)

125961-82-2 CAPLUS

Butanoic acid, 4-[6-acetyl-3-[3-[(4-acetyl-3-hydroxy-2-

propylphenyl)thio]propoxy]-2-propylphenoxy]- (9CI) (CA INDEX NAME)

$$HO_2C-(CH_2)_3-O$$
 AC
 $O-(CH_2)_3-S$
 $O-(CH_2)_3-S$

ANSWER 4 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1995:403614 CAPLUS

DOCUMENT NUMBER:

122:290448

TITLE:

Preparation of (acetylpropylphenoxy)alkanoic acids as

intermediates for antiallergic leukotriene antagonists INVENTOR(S):

Matsumoto, Toyomi; Aizawa, Yasuhiro; Matsukubo,

Hiroshi

PATENT ASSIGNEE(S):

Kyorin Seiyaku Kk, Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 3 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

PRI

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|----------------------|--------|------------|-----------------|------------|
| | | | | |
| JP 06345682 | A2 | 19941220 | JP 1993-166354 | 19930611 < |
| IORITY APPLN. INFO.: | | | JP 1993-166354 | 19930611 |
| HER SOURCE(S): | MARPAT | 122.290448 | | |

OTH GI

Y (CH₂)
$$_{m}X$$
 — COMe

Pr O (CH₂) $_{n}CO_{2}H$

AB The title compds. I (m = 2-5; n = 3-8; X = 0, S, S0, S02; Y = halo) are claimed. An aqueous NaOH solution was added dropwise to an EtOH solution of 4-[6-acetyl-3-(3-chloropropoxy)-2-propylphenoxy]butyric acid Et ester (preparation given) at 18-28° and the reaction mixture was stirred at room temperature for 2 h to give 91% 4-[6-acetyl-3-(3-chloropropoxy)-2propylphenoxy]butyric acid (II). II (21.4 g) and 15.1 g 2-hydroxy-4-mercapto-3-propylacetophenone were dissolved in DMF and the solution was treated with K2CO3 under stirring at room temperature for 3 h to give 24.4 g 4-[6-acetyl-3-[3-(4-acetyl-3-hydroxy-2-propylphenylthio)propoxy]-2propylphenoxy]butyric acid as a leukotriene antagonist.

125961-82-2P IT

RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)

(preparation of (acetylpropylphenoxy) alkanoic acids as intermediates for leukotriene antagonists)

L4 ANSWER 5 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1995:39068 CAPLUS

DOCUMENT NUMBER: 123:169347

TITLE: preparation of phenylthiopropoxyphenyloxybutyric acid

derivatives as leukotriene antagonists

INVENTOR(S): Oohashi, Mitsuo; Hori, Wataru

PATENT ASSIGNEE(S): Kyorin Seiyaku Kk, Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 13 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE | | |
|------------------------|--------|------------|-----------------|------------|--|--|
| | | | | · | | |
| JP 06100526 | A2 | 19940412 | JP 1992-273717 | 19920917 < | | |
| PRIORITY APPLN. INFO.: | | | JP 1992-273717 | 19920917 | | |
| OTHER COIDER (C). | ****** | 100 160010 | | | | |

OTHER SOURCE(S): MARPAT 123:169347

AB Title derivs. I (A, B = CO, hydroxymethylene; E = H, OH, acetoxy; G, L = Et, acetyl, 1-hydroxyethyl, 2-hydroxyethyl, hydroxycarbonylmethyl, lower alkoxycarbonylmethyl; X = void, O, O2; R1 = H, lower alkyl; X = O, O2 and B = hydroxymethylene when A = carbonyl, E = H, and G = L = Et) or their alkali salts, acting as strong antagonists for leukotrienes C4, D4, and E4 and useful for antiasthmatics, are prepared Thus, treating 2'-hydroxy-3'-(2-hydroxypropyl)-4'-mercaptoacetophenone (prepared in 6 steps from 3-allyl-2,4-dihydroxyacetophenone) with Et 4-[6-acetyl-3-(3-bromopropoxy)-2-propylphenoxylbutyrate gave I (A = B = CO, E = H, G = 1-hydroxyethyl, L = Et, R1 = Et, X = void).

IT 167211-62-3P 167211-73-6P 167211-79-2P

167211-83-8P 167211-94-1P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of phenylthiopropoxyphenyloxybutyric acid derivs. as leukotriene antagonists)

RN 167211-62-3 CAPLUS

CN Butanoic acid, 4-[6-acetyl-3-[3-[[4-acetyl-3-hydroxy-2-(2-hydroxypropyl)phenyl]thio]propoxy]-2-propylphenoxy]- (9CI) (CA INDEX NAME)

OH
$$CH_2-CH-Me$$

OH $CH_2-CH-Me$

RN 167211-73-6 CAPLUS

Butanoic acid, 4-[6-acetyl-3-[3-[[4-acetyl-3-hydroxy-2-(3-hydroxypropyl)phenyl]thio]propoxy]-2-propylphenoxy]- (9CI) (CA INDEX NAME)

 $HO_2C-(CH_2)_3-0$

CN

CN

RN 167211-79-2 CAPLUS

Butanoic acid, 4-[6-acetyl-3-[3-[(4-acetyl-3-hydroxy-2-propylphenyl)thio]propoxy]-2-(2-hydroxypropyl)phenoxy]- (9CI) (CA INDEX NAME)

O- (CH₂)₃-S

OH

$$CH_2$$
-CH-Me

 AC
 CH_2 -CH-Me

 AC
 AC
 CH_2 -CH-Me

RN 167211-83-8 CAPLUS

CN Butanoic acid, 4-[6-acetyl-3-[3-[(4-acetyl-3-hydroxy-2-propylphenyl)thio]propoxy]-2-(3-hydroxypropyl)phenoxy]- (9CI) (CA INDEX NAME)

O-
$$(CH_2)_3$$
 - S OH

AC $(CH_2)_3$ - OH

AC $(CH_2)_3$ - OH

RN 167211-94-1 CAPLUS

CN Butanoic acid, 4-[3-[3-[(4-acetyl-3-hydroxy-2-propylphenyl)thio]propoxy]-6-(hydroxyacetyl)-2-propylphenoxy]- (9CI) (CA INDEX NAME)

.
$$O-(CH_2)_3-S$$
 OH $O-(CH_2)_3-CO_2H$ $O-(CH_2)_3-CO_2H$

L4 ANSWER 6 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1990:138760 CAPLUS

DOCUMENT NUMBER: 112:138760

TITLE: Preparation of phenoxyalkylcarboxylic acid derivatives

as antiallergic agents.

INVENTOR(S): Ohashi, Mitsuo; Awano, Katsuya; Tanaka, Toshio;

Kimura, Tetsuya

PATENT ASSIGNEE(S): Kyorin Pharmaceutical Co., Ltd., Japan

SOURCE: Eur. Pat. Appl., 32 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GI

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE | |
|------------------------|------------|-----------|-----------------|-------------|--|
| | - - | | | | |
| • | A1 | 19890913 | EP 1989-103897 | 19890306 < | |
| EP 332109 | B1 | 19911204 | | | |
| R: BE, CH, DE, | ES, FR | , GB, IT, | LI, NL, SE | | |
| JP 02001459 | A2 | 19900105 | JP 1989-38912 · | 19890218 < | |
| JP 07116125 | B4 | 19951213 | | | |
| US 4985585 | A | 19910115 | US 1989-313900 | 19890223 < | |
| AU 8930884 | A1 | 19890907 | AU 1989-30884 | 19890301 < | |
| AU 617439 | B2 | 19911128 | | | |
| CA 1331763 | A1 | 19940830 | CA 1989-592555 | 19890302.< | |
| HU 50112 | A2 | 19891228 | HU 1989-1039 . | 19890303 < | |
| HU 204030 | В | 19911128 | | | |
| HU 208418 | В | 19931028 | HU 1991-2410 | 19890303 < | |
| HU 208524 | В | 19931129 | HU 1991-2411 | 19890303 < | |
| ES 2045219 | Т3 | 19940116 | ES 1989-103897 | 19890306 < | |
| CN 1036560 | Α | 19891025 | CN 1989-101301 | 19890307 < | |
| CN 1022407 | В | 19931013 | | | |
| PRIORITY APPLN. INFO.: | | | JP 1988-53374 | A 19880307 | |
| | | | | A3 19890303 | |
| OTHER SOURCE(S): | MARPAT | 112:13876 | | | |
| | | | | | |

MeCO
$$X^1 (CH_2)_m X^2$$
 COMe
HO Pr Pr O(CH₂)_nCO₂R¹ I

AB The title compds. (I; R1 = H, Me, Et; X1, X2 = O, S, SO, SO2; X1 = X2 \neq O; m = 2-5; n = 3-8), useful as antiallergic agents, are prepared A mixture of phenoxybutyrate II, bromopropyl thioether III, KI, and K2CO3 in

Me2CO was refluxed to give 72.4% I (R1 = Et, X1 = S, X2 = O, m = n = 3). I showed 66.7-96.2% inhibition of leukotriene D4-induced bronchoconstriction at 50 mg/kg p.o. in guinea pigs. Addnl. 70 I were also prepared

IT 125961-82-2P 125961-92-4P 125961-93-5P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of, as antiallergic agent)

RN 125961-82-2 CAPLUS

CN

RN

Butanoic acid, 4-[6-acetyl-3-[3-[(4-acetyl-3-hydroxy-2-propylphenyl)thio]propoxy]-2-propylphenoxy]- (9CI) (CA INDEX NAME)

125961-92-4 CAPLUS

CN Butanoic acid, 4-[6-acetyl-3-[3-[(4-acetyl-3-hydroxy-2-propylphenyl)sulfinyl]propoxy]-2-propylphenoxy]- (9CI) (CA INDEX NAME)

RN 125961-93-5 CAPLUS

CN Butanoic acid, 4-[6-acetyl-3-[3-[(4-acetyl-3-hydroxy-2-propylphenyl)sulfonyl]propoxy]-2-propylphenoxy]- (9CI) (CA INDEX NAME)

L4 ANSWER 7 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1983:575604 CAPLUS

DOCUMENT NUMBER: 99:175604

TITLE: Anti-SRS-A carboxylic acid derivatives and

pharmaceutical formulations containing them

INVENTOR(S): Bantick, John Raymond

PATENT ASSIGNEE(S): Fisons Ltd., UK

SOURCE: Eur. Pat. Appl., 67 pp.

CODEN: EPXXDW

Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

DOCUMENT TYPE:

PATENT NO.

KIND DATE

APPLICATION NO.

DATE

| • | | | | | |
|------------------------|--------|--------------|----------------|---|------------|
| | | | | - | |
| EP 79637 | A1 | 19830525 | EP 1982-201368 | | 19821101 < |
| EP 79637 | B1 | 19870128 | | | |
| R: AT, BE, CH, | DE, FR | , GB, IT, LI | , LU, NL, SE | | |
| US 4474788 | Α | 19841002 | US 1982-438163 | | 19821101 < |
| AT 25251 | E | 19870215 | AT 1982-201368 | | 19821101 < |
| JP 58090557 | A2 | 19830530 | JP 1982-196883 | | 19821111 < |
| PRIORITY APPLN. INFO.: | | | GB 1981-34186 | Α | 19811112 |
| | | | EP 1982-201368 | Α | 19821101 |
| CT. | | | | • | |

$$\begin{array}{c}
R \\
R_1 \\
R^2
\end{array}$$

$$\begin{array}{c}
R^5 \\
R^4
\end{array}$$

III

Ι

AB Anti-allergy (no data) bicyclic compds. I [R, R1 = H, alkyl; RR1 = bond; R2 = CO2H, carboxyalkyl; R3 = substituted OH, SH, NH2; R4, R5 = H, halogen, (un) substituted OH, NH2, alkyl, acyl; X = S, O, NR6 (R6 = H, alkyl)] were prepared Thus, 3,2,4-Pr(HO)2C6H2Ac reacted with 4,2,3-AcPr(H2N)C6H2S(CH2)3Br to give phenol II, which cyclized with EtO2CCO2Et to give quinoline III [R7 = Et, R8R9 = CH:C(CO2Et)]. The latter compound gave III (R7 = H, R8 = Me, R9 = H) on hydrolysis.

IT 87472-35-3P 87472-36-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 87472-35-3 CAPLUS

CN 4H-1-Benzopyran-2-propanoic acid, 7-[3-[(4-acetyl-3-hydroxy-2-propylphenyl)thio]propoxy]-4-oxo-8-propyl- (9CI) (CA INDEX NAME)

$$S-(CH_2)_3-O$$
 O
 $CH_2-CH_2-CO_2H$
 O
 O
 O

87472-36-4 CAPLUS

CN L-Lysine, mono[7-[3-[(4-acetyl-3-hydroxy-2-propylphenyl)thio]propoxy]-4-oxo-8-propyl-4H-1-benzopyran-2-propanoate] (9CI) (CA INDEX NAME)

CM 1

RN

CRN 87472-35-3 CMF C29 H34 O7 S

$$S-(CH_2)_3-O$$
 $CH_2-CH_2-CO_2H$
 O
 O

CM 2

CRN 56-87-1

CMF C6 H14 N2 O2

Absolute stereochemistry.